=> fil reg; s q.yn.pnptad.ktav..ssdf.a.li/sqsp FILE 'REGISTRY' ENTERED AT 10:32:06 ON 10 JUN 94 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 1994 American Chemical Society (ACS)

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L1 16 Q.YN.PNPTAD.KTAV..SSDF.A.LI/SQSP

=> d 1-16 .bevreg; fil ca; s 11

L1 ANSWER 1 OF 16 REGISTRY COPYRIGHT 1994 ACS

RN 152479-24-8 REGISTRY

CN INDEX NAME NOT YET ASSIGNED

SQL 77

MF Unspecified

CI MAN

L1 ANSWER 2 OF 16 REGISTRY COPYRIGHT 1994 ACS

RN 133723-39-4 REGISTRY

CN Glycolipoprotein MACIF (human clone p352-3 protein moiety reduced), N-L-methionyl- (9CI) (CA INDEX NAME)

SQL 104

MF Unspecified

CI MAN

L1 ANSWER 3 OF 16 REGISTRY COPYRIGHT 1994 ACS

RN 133722-42-6 REGISTRY

CN 1-86-Antigen CD 59 (human clone p352-3 protein moiety reduced), N-L-methionyl- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 1-86-Glycolipoprotein MACIF (human clone p352-3 protein moiety reduced), N-L-methionyl-

SOL 87

MF C424 H654 N116 O149 S11

CI MAN

L1 ANSWER 4 OF 16 REGISTRY COPYRIGHT 1994 ACS

RN 133722-41-5 REGISTRY

CN 1-86-Antigen CD 59 (human clone p352-3 protein moiety reduced) (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 1-86-Glycolipoprotein MACIF (human clone p352-3 protein moiety reduced)

SOL 86

MF C419 H645 N115 O138 S10

CI MAN

L1 ANSWER 5 OF 16 REGISTRY COPYRIGHT 1994 ACS

RN 133722-40-4 REGISTRY

CN 1-82-Antigen CD 59 (human clone p352-3 protein moiety reduced), N-L-methionyl- (9CI) (CA INDEX NAME)

OTHER NAMES:

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OTHER NAMES:
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    . reduced), N-L-methionyl-
SQL
MF
     C406 H623 N111 O141 S11
CI
   MAN
                     REGISTRY COPYRIGHT 1994 ACS
L1
     ANSWER 6 OF 16
RN
     133722-39-1 REGISTRY
     1-82-Antigen CD 59 (human clone p352-3 protein moiety reduced) (9CI)
CN
     (CA INDEX NAME)
OTHER NAMES:
     1-82-Glycolipoprotein MACIF (human clone p352-3 protein moiety
CN
SQL
     82
MF
     C401 H614 N110 O130 S10
CI
     MAN
     ANSWER 7 OF 16 REGISTRY COPYRIGHT 1994 ACS
L1
     133722-36-8 REGISTRY
RN
     1-77-Antigen CD 59 (human clone p352-3 protein moiety reduced),
CN
     N-L-methionyl- (9CI) (CA INDEX NAME)
OTHER NAMES:
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CN
     reduced), N-L-methionyl-
SOL
     C389 H594 N106 O134 S11
MF
   MAN
CI
     ANSWER 8 OF 16 REGISTRY COPYRIGHT 1994 ACS
L1
RN
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CN
     N-L-methionyl- (9CI) (CA INDEX NAME)
OTHER NAMES:
     1-76-Glycolipoprotein MACIF (human clone p352-3 protein moiety
CN
     reduced), N-L-methionyl-
SQL
MF
     C385 H588 N104 O122 S11
CI
     MAN
     ANSWER 9 OF 16 REGISTRY COPYRIGHT 1994 ACS
L1
     133722-34-6 REGISTRY
RN
     1-77-Antigen CD 59 (human clone p352-3 protein moiety reduced) (9CI)
CN
     (CA INDEX NAME)
OTHER NAMES:
CN
     1-77-Glycolipoprotein MACIF (human clone p352-3 protein moiety
     reduced)
SQL
     77
MF
     C384 H585 N105 O123 S10
CI
     MAN
L1
     ANSWER 10 OF 16 REGISTRY COPYRIGHT 1994 ACS
RN
     133722-33-5 REGISTRY
CN
     1-75-Antigen CD 59 (human clone p352-3 protein moiety reduced),
     N-L-methionyl- (9CI) (CA INDEX NAME)
OTHER NAMES:
CN
     1-75-Glycolipoprotein MACIF (human clone p352-3 protein moiety
     reduced), N-L-methionyl-
SQL
     76
MF
     C380 H581 N103 O129 S11
CI
     MAN
```

1 .

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ANSWER 11 OF 16 REGISTRY COPYRIGHT 1994 ACS
L1
RN
    133722-32-4 REGISTRY
     1-76-Antigen CD 59 (human clone p352-3 protein moiety reduced) (9CI)
CN
      (CA INDEX NAME)
OTHER NAMES:
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CN
     reduced)
SQL
     76
MF
     C380 H579 N103 O121 S10
CI
     MAN
                      REGISTRY COPYRIGHT 1994 ACS
L1
     ANSWER 12 OF 16
RN
     133722-31-3 REGISTRY
     1-75-Antigen CD 59 (human clone p352-3 protein moiety reduced) (9CI)
CN
      (CA INDEX NAME)
OTHER NAMES:
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CN
SOL
     75
     C375 H572 N102 O118 S10
MF
CI
     MAN
     ANSWER 13 OF 16 REGISTRY COPYRIGHT 1994 ACS
L1
RN
     133722-30-2 REGISTRY
     1-70-Antigen CD 59 (human clone p352-3 protein moiety reduced),
CN
     N-L-methionyl- (9CI) (CA INDEX NAME)
OTHER NAMES:
     1-70-Glycolipoprotein MACIF (human clone p352-3 protein moiety
CN
     reduced), N-L-methionyl-
SQL
     71
MF
     C351 H540 N96 O120 S11
CI
     MAN
                                 COPYRIGHT 1994 ACS
L1
     ANSWER 14 OF 16
                      REGISTRY
RN
     133722-28-8 REGISTRY
     1-70-Antigen CD 59 (human clone p352-3 protein moiety reduced) (9CI)
CN
      (CA INDEX NAME)
OTHER NAMES:
     1-70-Glycolipoprotein MACIF (human clone p352-3 protein moiety
CN
     reduced)
SQL
     70
MF
     C346 H531 N95 O119 S10
CI
     MAN
     ANSWER 15 OF 16 REGISTRY COPYRIGHT 1994 ACS
L1
RN
     126546-13-2 REGISTRY
     Antigen CD 59 (human clone YTH 53.1/1 protein moiety reduced) (9CI)
CN
      (CA INDEX NAME)
OTHER NAMES:
CN
     Antigen 1F 5 (human protein moiety reduced)
     Antigen CD 59 (human clone K-562-3 protein moiety reduced)
CN
ĊN
     Antigen CD 59 (human clone R18 protein moiety reduced)
     Glycolipoprotein HRF 20 (human clone pUIF10 protein moiety reduced)
CN
CN
     Glycolipoprotein HRF 20 (human clone pUIF10 reduced)
     Glycolipoprotein MACIF (human clone p352-3 protein moiety reduced)
CN
SQL
     103
MF
     Unspecified
CI
     MAN
     ANSWER 16 OF 16 REGISTRY
                                 COPYRIGHT 1994 ACS
L1
     126546-12-1 REGISTRY
RN
```

44

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RN
     126546-12-1 REGISTRY
CN
     Antigen CD 59 (human clone YTH53.1/1 precursor protein moiety
   reduced) (9CI) (CA INDEX NAME)
OTHER NAMES:
     Antigen 1F 5 (human precursor protein moiety reduced)
CN
     Antigen CD 59 (human clone K-562-3 precursor protein moiety reduced)
CN
CN
     Antigen CD 59 (human clone R18 precursor protein moiety reduced)
     Glycolipoprotein HRF 20 (human clone pUIF10 precursor protein moiety
CN
     Glycolipoprotein HRF 20 (human clone pUIF10 precursor reduced)
CN
CN
     Glycolipoprotein MACIF (human clone p352-3 precursor protein moiety
SQL
     128
MF
     Unspecified
CI
    MAN
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FILE COVERS 1967 - 28 May 1994 (940528/ED) VOL 120 ISS 22
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  HCA File when conducting SmartSELECT searches with large
  numbers of terms.
L2
            13 L1
=> d 1-13 .bevstr; fil caprev; s l1
     ANSWER 1 OF 13 CA COPYRIGHT 1994 ACS
L2
AN
     CA120(7):71831d CA
TI
     Determination of carboxyl-terminal residue and disulfide bonds of
    MACIF (CD59), a glycosyl-phosphatidylinositol-anchored membrane
     Sugita, Yuji; Nakano, Yasuko; Oda, Eiichi; Noda, Keiichi; Tobe,
ΑU
     Takashi; Miura, Nam Ho; Tomita, Motowo
CS
     Sch. Pharm. Sci., Showa Univ.
LO
     Tokyo 142, Japan
SO
     J. Biochem. (Tokyo), 114(4), 473-7
SC
     6-3 (General Biochemistry)
DΤ
     J
CO
     JOBIAO
IS
     0021-924X
PY
     1993
LA
     Eng
AN
     CA120(7):71831d CA
AB
    MACIF (CD59) is a glycosyl-phosphatidylinositol (GPI)-anchored
    membrane glycoprotein which inhibits the formation of the membrane
     attack complex of human complement. MACIF prepd. from human
     erythrocyte membranes was digested with pronase. When the digest was
     subjected to 2-phase partition with BuOH and 0.1N HCl, the
     C-terminal peptide was recovered in the BuOH phase because of the
     attachment of the highly hydrophobic GPI. The amino acid sequence of
     the peptide was detd. to be Asn-72 at its N-terminus and up to
     Glu-76, whhereas the presence of Asn-77 was ambiguous. To allow
     unequivocal detn. of the C-terminus, a sol. form of MACIF was prepd.
     from human urine on a large scale. The C-terminal peptide from the
     sol. form was prepd. by tryptic digestion followed by reversed-phase
     HPLC. The sequence and compn. of the peptide unequivocally revealed
```

HPLC. The sequence and compn. of the peptide unequivocally revealed Asn-77 as the C-terminus. The pattern of disulfide bonds of MACIF was also detd. with the membrane form as well as the sol. form. Cystine-contg. peptides were prepd. by chymotryptic and tryptic digestion, purified by HPLC, and their amino acid sequences were detd. The results indicated that disulfide bonds were formed at Cys-3-Cys-26, Cys-6-Cys-13, Cys-19-Cys-39, Cys-45-Cys-63(or 64), and Cys-63 (or 64) -Cys-69. IT 152479-24-8 (of erythrocytes, of human) ANSWER 2 OF 13 CA COPYRIGHT 1994 ACS CA119(17):175211x CA Structure of the CD59-encoding gene: further evidence of a relationship to murine lymphocyte antigen Ly-6 protein. [Erratum to document cited in CA119(7):64326u] Petranka, John G.; Fleenor, Donald E.; Sykes, Kathryn; Kaufman, Russel E.; Rosse, Wendell F. Med. Cent., Duke Univ. Durham, NC 27710, USA Proc. Natl. Acad. Sci. U. S. A., 90(12), 5878 3-3 (Biochemical Genetics) 13, 15 J PNASA6 0027-8424 1993 Eng CA119(17):175211x CA An error in mapping a 4.5-kb EcoRI restriction band has been cor. The size est. for intron 1 has consequently been increased. A revised restriction map for intron 1 has been presented. The error was not reflected in the abstr. or the index entries. IT 126546-12-1, Antigen CD 59 (human clone YTH53.1/1 precursor protein moiety reduced) (amino acid sequence of, complete (Erratum)) CA COPYRIGHT 1994 ACS ANSWER 3 OF 13 CA119(7):64326u CA Structure of the CD59-encoding gene: Further evidence of a relationship to murine lymphocyte antigen Ly-6 protein Petranka, John G.; Fleenor, Donald E.; Sykes, Kathryn; Kaufman, Russel E.; Rosse, Wendell F. Med. Cent., Duke Univ. Durham, NC 27710, USA Proc. Natl. Acad. Sci. U. S. A., 89(17), 7876-9 3-3 (Biochemical Genetics) 13, 15 J PNASA6 0027-8424 1992 CA119(7):64326u CA The gene for CD59 [membrane inhibitor of reactive lysis (MIRL), protectin], a phosphatidylinositol-linked surface glycoprotein that regulates the formation of the polymeric C9 complex of complement and that is deficient on the abnormal hematopoietic cells of patients with paroxysmal nocturnal hemoglobinuria, consists of four exons spanning 20 kilobases. The untranslated first exon is preceded by a G+C-rich promoter region that lacks a consensus TATA or CAAT

motif. The second exon encodes the hydrophobic leader sequence of

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LO

SO

SC SX

DT

CO

IS

PY

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AB

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∌AU

CS LO

SO

SC

SX DT

.»CO

IS

PY

ĹΑ AN

AB

motif. The second exon encodes the hydrophobic leader sequence of the protein, and the third exon encodes the amino-terminal portion of the mature protein. The fourth exon encodes the remainder of the mature protein, including the hydrophobic sequence necessary for glycosyl-phosphatidylinositol anchor attachment. The structure of the CD59 gene is very similar to that encoding Ly-6, a murine glycoprotein with which CD59 has some structural similarity. The striking similarity in gene structure is further evidence that the two proteins belong to a superfamily of proteins that may also include the urokinase plasminogen-activator receptor and a squid glycoprotein of unknown function.

IT 126546-12-1, Antigen CD 59 (human clone YTH53.1/1 precursor protein moiety reduced)

(amino acid sequence of, complete)

```
L2 ANSWER 4 OF 13 CA COPYRIGHT 1994 ACS
```

AN CA118(20):198171c CA

TI Genetically engineered cells as universal donor cells for vascular grafts or drug delivery

IN Sims, Peter J.; Bothwell, Alfred L. M.; Elliot, Eileen A.; Flavell, Richard A.; Madri, Joseph; Rollins, Scott; Bell, Leonard; Squinto, Stephen

PA Oklahoma Medical Research Foundation; Yale University

LO USA

SO PCT Int. Appl., 88 pp.

PI WO 9302188 A1 930204

DS W: CA, JP

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, MC, NL, SE

AI WO 92-US5920 920714

PRAI US 91-729926 910715

US 92-906394 920629

SC 63-3 (Pharmaceuticals)

SX 3, 15

DT P

CO PIXXD2

PY 1993

LA Enq

AÑ CA118(20):198171c CA

Genetically engineered cells are provided which can serve as AB universal donor cells in such applications as reconstruction of vascular linings or the administration of therapeutic agents. The cells include a DNA sequence which is expressed by the cell and which codes for a protein having complement inhibitory activity and which provides protection against complement-based lysis, i.e., hyperacute rejection. In addn., the cell's natural genome is changed so that proteins encoded by either the class II or both the class I and the class II major histocompatibility complex genes do not appear on the cell's surface. In this way, attack by T-cells is avoided. Optionally, the cells can include a self-destruction mechanism so that they can be removed from the host when no longer needed. The cells may further comprise of polynucleotide coding for a therapeutic agent which is expressed and secreted by the cell. When cDNA encoding the human CD59 antigen was stably incorporated into the genome of porcine aortic endothelial cells (PAEC) and expressed on the cell surface, the cells were protected from complement-mediated attack as assayed by human complement-mediated cell lysis in vitro. The recombinant PAEC had similar biol. behavior as normal PAEC in terms of proliferation rates, not overgrowing monolayers or growing in suspension, and being contact inhibited. Addnl., the recombinant PAEC were capable of attaching to a synthetic Gortex graft as well as normal endothelial cells.

IT 126546-13-2, CD59 antigen (human reduced)

```
IT 126546-13-2, CD59 antigen (human reduced)
         (amino acid sequence of and expression of gene for, on cell
         surface of recombinant cells)
     ANSWER 5 OF 13 CA COPYRIGHT 1994 ACS
L2
AN
     CA115(17):176706s CA
     Molecular cloning of cDNA for human lymphocyte surface antigen CD59
ΤI
     Sawada, Ritsuko; Naruto, Masanobu
 IN
PA
     Toray Industries, Inc.
 LO
     Japan
 SO
     Jpn. Kokai Tokkyo Koho, 16 pp.
 PΙ
     JP 03081297 A2 910405 Heisei
     JP 89-218183 890823
ΑI
     3-4 (Biochemical Genetics)
 SC
 SX
 DT
 CO
     JKXXAF
 PY
     1991
 LA
     Japan
AN
     CA115(17):176706s CA
     The cDNA encoding human lymphocyte surface antigen CD59, a human
 AB
     counterpart of the mouse antigen Ly-6 and that is recognized by the
     anti-Ly-6 monoclonal antibody MEM43, is cloned and sequenced. A cDNA
      library prepd. from human peri[heral monocytes was screened using a
     350 base pair probe, which was obtained in a preparatory
     hybridization using 3 synthetic oligonucleotide probes (Ly-61, Ly-2,
     and Ly-3) that encoded the N-terminal amino acids of an antigen
     recognized by MEM43. Clones 5 and 18 contg. the same CD59-encoding
     cDNA but different flanking regions were isolated.
 IT 126546-12-1, Antigen CD 59 (human clone YTH53.1/1 precursor
     protein moiety reduced) 126546-13-2, Antigen CD 59 (human
     clone YTH 53.1/1 protein moiety reduced)
         (amino acid sequence of and cloning in Escherichia coli of cDNA
         for)
 L2
     ANSWER 6 OF 13 CA
                         COPYRIGHT 1994 ACS
 AN
      CA115(7):66207x CA
     Human lymphocyte surface antigen comparable to mouse Ly6 antigen and
 TΙ
      its cDNA cloning
 IN
      Sawada, Ritsuko; Naruto, Masanobu
 PA
     Toray Industries, Inc.
 LO
      Japan
 SO
     Jpn. Kokai Tokkyo Koho, 12 pp.
      JP 03048696 A2 910301 Heisei
 PI
AI
     JP 89-183264 890714
 SC
      3-4 (Biochemical Genetics)
 SX
     15
DT
     JKXXAF
 CO
 PY
      1991
 LA
     Japan
AN
      CA115(7):66207x CA
     A cDNA encoding a human lymphocyte surface antigen that is
AB
      comparable to the mouse lymphocyte antigen Ly6 is cloned, sequenced,
      and its amino acid sequence deduced. Three synthetic oligonucleotide
     probes were prepd. according to the 17 N-terminal amino acid
      sequences of human antigen CD59 that was recognized by monoclonal
      antibody MEM43 and was considered comparable to the Ly6 antigen. The
      cDNA libraries of human monocytic leukemic cell line J111 and
      peripheral lymphocytes were amplified by polymerase chain reaction
      and screened with the 3 probes to obtain clone P-1 carrying the cDNA
```

for the human counterpart of the mouse Ly6 antigen.

```
for the human counterpart of the mouse Ly6 antigen.
IT 126546-13-2, Antigen CD 59 (human clone YTH 53.1/1 protein
   moiety reduced)
        (amino acid sequence of and cloning in Escherichia coli of cDNA
    ANSWER 7 OF 13 CA COPYRIGHT 1994 ACS
L2
     CA114(21):201135u CA
AN
     Cloning and expression of human membrane attack complex inhibition
TI
     factor (MACIF) gene
     Tomita, Motowo; Sugita, Yuji; Takemoto, Toshiyuki; Furuichi,
IN
    Kiyoshi; Takayama, Makoto; Yusakawa, Ko; Yano, Shinya; Yamaji,
    Noboru; Ito, Katsuhisa
    Yamanouchi Pharmaceutical Co., Ltd.
PA
LO
    Japan
SO
     Eur. Pat. Appl., 49 pp.
PΙ
     EP 394035 A2 901024
       AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE
DS
    R:
ΑI
     EP 90-304203 900419
PRAI JP 89-103088 890421
    JP 89-179933 890712
     JP 89-230983 890906
    JP 89-238246 890913
    JP 89-247818 890921
    JP 89-281197 891027
SC
     3-4 (Biochemical Genetics)
DT
    P
CO
    EPXXDW
PY
    1990
LA
    Enq
os
    MARPAT 114:201135
     CA114(21):201135u CA
AN
     Human MACIF (I) and its biol. active fragment cDNAs are cloned and
AB
     expressed in animal or microbial cells. From a cDNA library of human
     monocyte in pGEM4, I cDNA was cloned by the colony hybridization
    method using 2 DNA probes encoding N and C-terminal fragments,
     resp., of I. I was produced in CHO cells as a phosphatidylinositol-
     anchored protein on the cell membrane. Also given was the expression
     of cDNA for I and its fragments in Escherichia coli and CHO cells.
IT 133722-30-2 133722-33-5 133722-35-7
   133722-36-8 133722-40-4 133722-42-6
   133723-39-4
        (amino acid sequence of)
IT 126546-12-1, Antigen CD 59 (human clone YTH53.1/1 precursor
     protein moiety reduced)
        (amino acid sequence of and cloning in Escherichia coli of cDNA
        for)
IT 133722-28-8, 1-70-Antigen CD 59 (human clone p352-3 protein
     moiety reduced) 133722-31-3, 1-75-Antigen CD 59 (human
     clone p352-3 protein moiety reduced) 133722-32-4,
     1-76-Antigen CD 59 (human clone p352-3 protein moiety reduced)
   133722-34-6, 1-77-Antigen CD 59 (human clone p352-3 protein
     moiety reduced) 133722-39-1, 1-82-Antigen CD 59 (human
     clone p352-3 protein moiety reduced) 133722-41-5,
     1-86-Antigen CD 59 (human clone p352-3 protein moiety reduced)
        (amino acid sequence of and expression in CHO cells of cDNA for)
     ANSWER 8 OF 13 CA COPYRIGHT 1994 ACS
L2
AN
     CA113(19):166413p CA
     Isolation and expression of the full-length cDNA encoding CD59
TI
     antigen of human lymphocytes
     Sawada, Ritsuko; Ohashi, Kensaku; Anaguchi, Hiroyuki; Okazaki,
ΑU
```

```
Sawada, Ritsuko; Ohashi, Kensaku; Anaguchi, Hiroyuki; Okazaki,
AU
     Hitoaki; Hattori, Masakazu; Minato, Nagahiro; Naruto, Masanobu
    · Basic Res. Lab., Toray Ind., Inc.
CS
LO
     Kamakura 248, Japan
     DNA Cell Biol., 9(3), 213-20
SO
SC
     3-3 (Biochemical Genetics)
SX
     13, 15
DT
     J
CO
     DCEBE8
IS
     1044-5498
PY
     1990
LA
     Eng
AN
     CA113(19):166413p CA
     To identify the primary structure of CD59 antigen and to elucidate
AB
     its function, a full-length cDNA clone of CD59 was isolated. The
     cDNA sequence contained an open reading frame that encodes a
     128-amino-acid peptide. The amino-terminal 25 amino acids
     represented a typical signal peptide sequence, and the
     carboxy-terminal hydrophobic amino acids were characteristic for
     phosphatidylinositol-anchored proteins. The predicted mature protein
     sequence showed 35% homol. with murine Ly-6C.1 and 31% with Ly-6A.2.
     The no. and the distribution of cysteine residues were conserved,
     implying that the CD59 represented a human homolog of murine Ly-6.
     RNA blot hybridization anal. revealed the expression of CD59 mRNA in
     placental, lung, and pancreatic tissues. The mRNA was not only
     expressed in T-cell lines but in some monocytic, myeloid, and B-cell
     lines. In all of these tissues and cell lines, at least 4 mRNA
     species were detected. DNA blot hybridization anal. revealed a
     rather simple genomic structure, which suggested a single gene as
     compared with the complex multigene family of murine Ly-6.
IT 129817-69-2, Antigen CD 59 (human clone R18 precursor
     protein moiety reduced) 129817-70-5, Antigen CD 59 (human
     clone R18 protein moiety reduced)
        (amino acid sequence of)
L2
     ANSWER 9 OF 13
                     CA COPYRIGHT 1994 ACS
AN
     CA113(11):92818a CA
     Cloning of the gene for a human complement-mediated cell membrane
TI
     damage-inhibiting glycoprotein
     Okada, Hidechika; Okada, Noriko; Nagami, Yoichi; Takahashi,
IN
     Kazuhiro; Takizawa, Hisao; Kondo, Jun
PA
     Mitsubishi Kasei Corp.
LO
     Japan
SO
     Eur. Pat. Appl., 25 pp.
PΙ
     EP 351313 A2 900117
DS
         BE, CH, DE, FR, GB, IT, LI, NL, SE
ΑI
     EP 89-401996 890711
PRAI JP 88-172187
                   880711
     JP 89-129944
                   890523
SC
     3-4 (Biochemical Genetics)
ŞX
     13, 15
ĎΤ
     P
CO
     EPXXDW
\mathbf{P}\mathbf{Y}
     1990
ĿΆ
     Eng
AN
     CA113(11):92818a
                       CA
*AB
     The gene for a complement-mediated cell-membrane damage-inhibiting
     N-glycosidated protein (20-25 kilodalton) contg.
     phosphatidylinositol was isolated from a human placental cDNA
     library in .lambda. gt11 using probes derived fromthe amino acid
     sequence of the antigen (antigen 1F5). The 1F5 antigen, recognized
     by antibody 1F5 isolated from a hybridoma that causes
```

by antibody 1F5 isolated from a hybridoma that causes complement-mediated hemolysis of neuraminidase-digested human erythrocytes, was isolated from solubilized human erythrocyte membranes. The 1F5 antigen was immune-affinity purified using 1F5 antibody and shownto contain phosphatidylinositol, was N-glycosidated, and had a mol. wt. of 20-25.times. 103. Poly (A)+ RNA of K562 cells was used to prep. 1F5 antigen cDNA probes by polymerase chain reaction, and the probes used to screen the cDNA library. Clones contq. 1.2-1.8 kb inserts were shown to contain common 500 bp Eco RI-Bam HI restriction fragment. Four clones contd. a full length cDNA for 1F5 antigen of 387 bp corresponding to a 128 amino acid protein including a 25 amino acid N-terminal signal peptide.

IT 128794-22-9, Antigen 1F 5 (human protein moiety reduced) (amino acid sequence of)

IT 128794-21-8, Antigen 1F 5 (human precursor protein moiety reduced)

(amino acid sequence of and cloning in Escherichia coli of cDNA for)

ANSWER 10 OF 13 CA COPYRIGHT 1994 ACS L2

CA113(7):56944e CA AN

The CD59 antigen is a structural homolog of murine Ly-6 antigens but TI lacks interferon inducibility

Philbrick, William M.; Palfree, Roger G. E.; Maher, Stephen E.; AU Bridgett, Margot M.; Sirlin, Sonia; Bothwell, Alfred L. M.

CS Med. Sch., Yale Univ.

LO New Haven, CT, USA

Eur. J. Immunol., 20(1), 87-92 SO

SC 15-2 (Immunochemistry)

DTJ

CO **EJIMAF**

IS 0014-2980

PΥ 1990

*LA

L2

AN CA113(7):56944e CA

A cDNA encoding the human leukocyte antigen CD59 has been isolated AB from the erythroid cell line K-562 and its identity confirmed through expression in COS cells. Northern blotting reveals 3 message species of approx. 800, 1400, and 2000 bases in size, which are constitutively expressed in all lymphoid, erythroid, myeloid, and neural cell types tested thus far. Southern blotting of human DNA indicates a pattern consistent with the presence of a single gene, which has been mapped to chromosome 11 by somatic cell hybrids. Also, the finding of a transcriptionally active cross-hybridizing gene in monkey cells suggests conservation of CD59 sequences among primates. Comparison of the CD59 protein sequence with those of the Ly-6E and Ly-6C antigens discloses a similarity in overall structure, including the alignment of abundant cysteine residues, hydrophobic C termini and conservation of amino acids surrounding the proposed phosphatidylinositol-glycan modification site for Ly-6 mols. Unlike Ly-6, however, CD59 expression does not appear to be inducible with interferons. This, along with its limited homol. and different tissue distribution, cast doubt upon the functional equivalence of CD59 and either of the well-characterized mouse Ly-6 proteins.

IT 128415-65-6, Antigen CD 59 (human clone K-562-3 precursor protein moiety reduced) 128415-66-7, Antigen CD 59 (human clone K-562-3 protein moiety reduced) (amino acid sequence of)

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L2
     ANSWER 11 OF 13 CA. COPYRIGHT 1994 ACS
AN
     CA112(21):196393d CA
ΤI
     20 KDa homologous restriction factor of complement resembles T cell
     activating protein
ΑU
     Okada, Hidechika; Nagami, Yoichi; Takahashi, Kazuhiro; Okada,
     Noriko; Hideshima, Teru; Takizawa, Hisao; Kondo, Jun
CS
     Sch. Med., Nagoya City Univ.
LO
     Nagoya 467, Japan
SO
     Biochem. Biophys. Res. Commun., 162(3), 1553-9
SC
     15-4 (Immunochemistry)
SX
DT
     J
CO
     BBRCA9
IS
     0006-291X
PY.
     1989
ĽΆ
     Eng
*AN
     CA112(21):196393d CA
AB
     The authors have previously identified a 20 KDa membrane
     glycoprotein 1F5 antigen which inhibits the assembly of homologous
     complement membrane attack complexes and it was designated as HRF20
     standing for 20 KDa homologous restriction factor. The amino acid
     sequence deduced from its coding base sequence resembles that of T
     cell-activating protein, most conspicuously in cysteine residues, 10
     out of 11 of which occupy identical positions in an overall sequence
     homol. of 24.8%. Proliferation of human T cells was stimulated by
     monoclonal antibody to HRF20.
IT 126805-73-0, Glycolipoprotein HRF 20 (human clone pUIF10
     precursor reduced) 126805-74-1, Glycolipoprotein HRF 20
     (human clone pUIF10 reduced)
        (amino acid sequence of)
     ANSWER 12 OF 13 CA COPYRIGHT 1994 ACS
L2
AN
     CA112(21):192901v CA
     Molecular cloning and characterization of MACIF, an inhibitor of
ΤĮ
     membrane channel formation of complement
     Sugita, Yuji; Tobe, Takashi; Oda, Eiichi; Tomita, Motowo; Yasukawa,
AU
     Ko; Yamaji, Noboru; Takemoto, Toshiyuki; Furuichi, Kiyoshi;
     Takayama, Makoto; Yano, Shinya
CS
     Sch. Pharm. Sci., Showa Univ.
ĽΟ
     Tokyo 142, Japan
SO
     J. Biochem. (Tokyo), 106(4), 555-7
SC
     3-3 (Biochemical Genetics)
SX
     6, 13
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CO
     JOBIAO
.IS
     0021-924X
PY
     1989
LA
     Eng
ÀΝ
     CA112(21):192901V CA
     Human erythrocytes contain a membrane protein, MACIF, which inhibits
AB
     the formation of a membrane attack complex (MAC) of complement. The
     authors cloned and sequenced the cDNA of MACIF mRNA. The amino acid
     sequence predicted from its nucleotide sequence consists of 128
     amino acids. The amino-terminal 25 residues may correspond to a
     signal peptide. The carboxy-terminal sequence confirmed that MACIF
     is a glycosylphosphatidylinositol (GPI)-anchored protein. The amino
     acid sequence of MACIF was partially detd. by established techniques
     for protein chem., and the resultant sequence was consistent with
     that predicted from the nucleotide sequence. The results of sequence
     analyses also suggested that asparagine at the 18th position was
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N-glycosylated. When mRNA obtained from the MACIF cDNA clone with SP6 RNA polymerase was microinjected into Xenopus oocytes, the

SP6 RNA polymerase was microinjected into Xenopus oocytes, the oocytes synthesized a product which exhibited MACIF activity and reacted with anti-MACIF antibody. Comparison of the predicted sequence revealed significant homol. with mouse Ly-6 antigens.

- IT 126880-43-1, Glycolipoprotein MACIF (human clone p352-3 precursor protein moiety reduced) 126880-44-2, Glycolipoprotein MACIF (human clone p352-3 protein moiety reduced) (amino acid sequence of)
- L2 ANSWER 13 OF 13 CA COPYRIGHT 1994 ACS
- AN CA112(19):176532V CA
- TI CD59, an LY-6-like protein expressed in human lymphoid cells, regulates the action of the complement membrane attack complex on homologous cells
- AU Davies, Alexandra; Simmons, David L.; Hale, Geoff; Harrison, Richard A.; Tighe, Helen; Lachmann, Peter J.; Waldmann, Herman
- CS Mol. Immunopathol. Unit, MRC Cent.
- LO Cambridge CB2 2QH, UK
- SO J. Exp. Med., 170(3), 637-54
- SC 15-2 (Immunochemistry)
- DT J
- CO JEMEAV
- IS 0022-1007
- PY 1989
- LA Eng
- AN CA112(19):176532V CA
- A novel cell surface antigen has been identified on a wide range of AB lymphoid cells and erythrocytes. A monoclonal antibody (mAb) YTH 53.1 (CD59) against this antigen enhanced the lysis of human red cells and lymphocytes by homologous complement. Studies of reactive lysis using different species of C.hivin.5.hivin.6, and of whole serum used as a source of C7-9, indicated that the inhibitory activity of the CD59 antigen is directed towards the homologous membrane attack complex. CD59 antigen was purified from human urine and erythrocyte stroma by affinity chromatog. using the mAb YTH 53.1 immobilized on Sepharose, and, following transient expression of a human T cell cDNA library in COS cells, the corresponding cDNA also identified using the antibody. The CD59 antigen is a small protein (.apprx.20 kD as judged by SDS-PAGE, 11.5 kD predicted from the isolated cDNA) sometimes assocd. with larger components (45 and 80 kD) in urine. The sequence of CD59 antigen is unlike that of other complement components or regulatory proteins, but shows 26% identity with that of the murine LY-6 antigen. CD59 antigen was released from the surface of transfected COS cells by phosphatidylinositolspecific phospholipase C, demonstrating that it is attached to the cell membrane by means of a glycolipid anchor; it is therefore likely to be absent from the surface of affected erythrocytes in the disease paroxysmal nocturnal hemoglobinuria.
- IT 126546-12-1, Antigen CD 59 (human clone YTH53.1/1 precursor protein moiety reduced) 126546-13-2, Antigen CD 59 (human clone YTH 53.1/1 protein moiety reduced) (amino acid sequence of)

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